Claims

 A method for classification of cancer in an individual having contracted cancer comprising

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 i) in a sample from the individual having contracted cancer determining the microsatellite status of the tumor and

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- ii) in a sample from the individual having contracted cancer, said sample comprising a plurality of gene expression products the presence and/or amount which forms a pattern, determining from said pattern a prognostic marker, wherein the microsatellite status and the prognostic marker is determined simultaneously or sequentially
- iii) classifying said cancer from the microsatellite status and the prognostic marker.

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The method according to claim 1, wherein the prognostic marker is the hereditary or sporadic nature of said cancer the determination of which comprises the steps of

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 i) in a sample from the individual having contracted cancer, said sample comprising a plurality of gene expression products the presence and/or amount of which forms a pattern that is indicative of the hereditary or sporadic nature of said cancer

ii) determining the presence and/or amount of said gene expression products forming said pattern,

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iii) obtaining an indication of the hereditary or sporadic nature of said cancer in the individual based on step ii).

The method of claims 1 or 2, wherein the determination of microsatellite status comprises the steps of

- i) in a sample from the individual having contracted cancer, said sample comprising a plurality of gene expression products the presence and/or amount of which forms a pattern that is indicative of the microsatellite status of said cancer,
- determining the presence and/or amount of said gene expression products forming said pattern,

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- iii) obtaining an indication of the microsatellite status of said cancer in the individual based on step ii).
- The method according to claims 1, 2 or 3, wherein the cancer is colon cancer.
- 5. The method of any of the preceding claims, wherein a plurality of gene expression products are analysed using solid support, having binding partners (hybridisation partners) for said plurality of gene expression products forming a pattern.
- The method of any of the preceding claims, wherein a plurality of gene expression products are analysed using binding partners (hybridisation partners) for said plurality of gene expression products forming a pattern.
- 7. The method of claims 1,2 or 3, wherein at least two of said plurality of gene expression products forming a pattern are used to determine said microsatellite status are selected individually from a group of genes indicative of microsatellite status.
- 8. The method of claims 1, 2 or 3, wherein at least two of said plurality of gene expression products used to determine the hereditary or sporadic nature of said colon cancer are selected individually from a group of genes indicative for the hereditary or sporadic nature of the cancer.
- The method of claims 1, 2 or 3, wherein at least two of said plurality of gene expression products forming a pattern used to determine said microsatellite status are selected individually from the group consisting of the genes listed below

Gene name	Ref seq	Gene symbol	SEQ ID NO.:
chemokine (C-C motif) ligand 5 Tryptophanyl-tRNA synthetase Proteasome (prosome, macropain) activa- tor subunit 1 (PA28 alpha)	NM_002985 NM_004184 NM_006263	CCL5 WARS PSME1	1 2 3
Bone marrow stromal cell antigen 2 ubiquitin-conjugating enzyme E2L 6	NM_004335 NM_004223	BST2 UBE2L6	4 5

			_
A kinase (PRKA) anchor protein 1	NM_003488	AKAP1	6
Proteasome (prosome, macropain) activa-	NM 002818	PSME2	7
tor subunit 2 (PA28 beta)			_
carcinoembryonic antigen-related cell	<u>NM_004363</u>	CEACAM5	8
adhesion molecule 5			
FERM, RhoGEF (ARHGEF) and pleck-		E1004	9
strin domain protein 1 (chondrocyte-	NM_005766	FARP1	
derived)			10
myosin X	NM_012334	MYO10	11
heterogeneous nuclear ribonucleoprotein	NIN 004500	LINDO	
L	NM_001533	HNRPL	12
Autocrine motility factor receptor	NM_001144	AMFR	13
dimethylargini ne dimethylaminohydrolase	NM 013974	DDAH2	
2	14W 010374		14
tumor necrosis factor, alpha-induced pro-	NM 006291	TNFAIP2	
tein 2 mutL homologi 1, colon cancer, nonpoly-	NM 000249	MLH1	15
posis type 2 (E. coli)	1011		
thymidylate synthetase	NM 001071	TYMS	16
intercellular adhesion molecule 1 (CD54),	NM 000201	ICAM1	17
human rhinovirus receptor			
general transcription factor IIA, 2, 12kDa	NM 004492	GTF2A2	18
Rho-associated, coiled-coil containing	NM 004850	ROCK2	19
protein kinase 2			
ATP binding protein associated with cell	NM_005783	TXNDC9	20
differentiation			0.4
NCK adaptor protein 2	NM_003581	NCK2	21
phytanoyl-CoA hydroxylase (Refsum dis-		-1001	22
ease)	NM 006214	PHYH	23
metastais-associated gene family, mem-	NIA 004720	MTA2	23
ber 2	NM 004739 NM 001091	ABP1	24
amiloride binding protein 1 (amine oxi-	MINI_001091	ADFI	<u>_</u>
dase (copper-containing))	NM: 000712	BLVRA	25
Biliverdin reductase A	NM 000933	PLCB4	26
phospholipase C, beta 4	NM 002416	CXCL9	27
chemokine (C-X-C motif) ligand 9		PURA	28
purine-rich element binding protein A	NM 005859 NM 014298	QPRT	29
quinolinate phosphoribosyltransferase	NW 014230	GI ICI	
(nicotinate-nucleotide pyrophosphorylase			
(carboxylating)) retinoic acid receptor responder (tazaro-	NM 004585	RARRES3	30
tene induced) 3			
chemokine (C-C motif) ligand 4	NM 002984	CCL4	31
forkhead box O3A	NM 001455	FOXO3A	32
interferon, alpha-inducible protein (clone	NM 002038	G1P3	34
IFI-6-16)	NM 022873		123
chemokine (C-X-C motif) ligand 10	NM 001565	CXCL10	35
diemokile (O-X-O moiii) iigano 10	NM 005950		36
metallothionein 1G	NM 005950		
	NM 000043		37
tumor necros is factor receptor super-	NM_152877		133
family, member 6	NM 152876		132
monipol o	NM 152875		134
•	NM_152872		130
	NM_152873		33 129
	NM_152871		129

NM_152874

to the transfer of the total of			38
endothelial cell growth factor 1 (platelet- derived)	NM 001953	ECGF1	30
SCO cytochrome oxidase deficient ho-	NM 005138	SCO2	39
molog 2 (yeast)	NIN 000440	0)(0) 40	40
chemokine (C-X-C motif) ligand 13 (B-cell chemoattractant)	NM 006419	CXCL13	40
Granulysin	NM_006433	GNLY	41
	_		
CD2 antigen (p50), sheep red blood cell			42
receptor	NM 001767	CD2	
splicing factor, arginine/serine-rich 6	NM_006275	SFRS6	43
Teratocarcinoma-derived growth factor 1	NM_003212	TDGF1	44
metallothionein 1H	NM 005951	MT1H	45
cytochrome P450, family 2, subfamily B,	NM_000767	CYP2B6	46
polypeptide 6			
tumor necrosis factor (ligand) superfamily,	NM_003811	TNFSF9	47
member 9	NIM 000047	RBM12	48
mana a sa	NM_006047	RBIVITZ	40
RNA binding motif protein 12	NM_006047 NM_006644	HSPH1	49
heat shock 105kDa/110kDa protein 1	NW 000044	поги	70
staufen. RNA binding protein (Drosophila)	NM 004602	STAU	50
Station, 144 (billiang proton) (Brosophine)	NM 017452		125
	NM_017453	•	126
I I I I I I I I I I I I I I I I I I I	NIM 021246	LY6G6D	51
lymphocyte antigen 6 complex, locus G6D	NM 021246 NM 007236	CHP	52
calcium binding protein P22		CDC14B	53
CDC14 cell division cycle 14 homolog B	NM_003671 NM_033331	CDC 14B	115
(S. cerevisiae)	MM_033331		110
Epiplakin 1	XM 372063	EPPK1	54
metallothionein 1X	NM 005952	MT1X	55
Transforming growth factor, beta receptor	NM 003242	TGFBR2	56
II (70/80kDa)			
protein kinase C binding protein 1	NM_012408	PRKCBP1	57
•	NM_183047		124
	**** 000070	TMACEC	58
Transmembrane 4 superfamily member 6	NM 003270	TM4SF6 PLEKHB1	59 ·
pleckstrin homology domain containing,	NM 021200	PLENTOI	28
family B (evectins) member 1	NIM ODDEC4	APOL1	60
apolipoprotein L, 1	NM_003661 NM_145343	W-OF1	120
	140040		3
Indoleamine-pyrrole 2,3 dioxygenase	NM 002164	INDO .	61
inducan inc-pyriole 2,0 dioxygonase	14141 002 104		

forkhead box A2	NM_021784	FOXA2	62
granzyme H (cathepsin G-like 2, protein h-CCPX)	NM 033423	GZMH	63
baculoviral IAP repeat-containing 3	NM_001165	BIRC3	64
Homo sapiens metallothionein 1H-like protein		AF333388 (Hs 382039)	135
KIAA0182 protein	NM 014615	KIAA0182	117
G protèin-coupled receptor 56	NM_005682	GPR56	65 116
metallothionein 2A	NM 201524 NM 005953	MT2A	66
F-box only protein 21	NM_015002	FBXO21	67
erythrocyte membrane protein band 4.1-like 1	NM_012156, NM_012156	EPB41L1	68
hypothetical protein MGC21416	NM 173834	MGC21416	69
protein O-fucosyltransferase 1	NM 015352.	POFUT1	70
proton o races jumanerales t	NM 015352		
metallothionein 1E (functional)	NM 175617	MT1E	71
troponin T1, skeletal, slow	NM 003283	TNNT1	72
chimerin (chimaerin) 2	NM 004067	CHN2	73
heterogeneous nuclear ribonu cleoprotein	1111_00 1001	0,	74
H1 (H)	NM 005520	HNRPH1	
ATP synthase, H+ transporting, mito- chondrial F1 complex, alpha subunit, iso- form 1, cardiac muscle	NM 004046	ATP5A1	75
eukaryotic translation initiation factor 5A	NM 001970	EIF5A	76
perforin 1 (pore forming protein)	NM 005041	PRF1	77
OGT(O-Glc-NAc transferase)-interacting	NM 014965	OIP106	78
protein 106 KDa			
DEAD (Asp-Glu-Ala-Asp) box polypeptide			79
27	NM_017895	DDX27	80
vacuolar protein sorting 35 (yeast)	NM_018206	VPS35	81
tripartite motif-containing 44	NM_017583	TRIM44	01
transmembrane, prostate androgen in- duced	NM 020182	TMEPAI	82
RNA	NM_199169	TIVILEPAI	127
1371	NM 199170		128
dynein, cytoplasmic, light polypeptide 2A	NM_014183	DNCL2A	83
	NM 177953		122
leucine aminopeptidase 3	NM 015907	LAP3	84
Chromosome 20 open reading frame 35	NM_018478	C20orf35	85
			-
	NM_033542		118
solute carrier family 38, member 1	NM_030674	SLC38A1	86

ATPase, class II, type 9a	9	ATP9a	
	<u>Xm 030577.</u>		104
associated)	1		
hypothetical protein FLJ20618 SET translocation (myeloid leukaemia-	NM 003011.		103
mitochondrial solute carrier protein	NM_016612 NM_017903	MSCP FLJ20618	101
. N	NINA 016610	MECD	101
	NM_145343		120
apolipoprotein L, 2	NM 030882,	APOL2	100
hypothetical protein FLJ20232	NM 019008	FLJ20232	99
	NM 138932		119
apobec-1 complementation factor	NM_014576,	ACF	98
aryl hydrocarbon receptor nuclear translo- cator-like 2	NM_020183	ARNTL2	-
NAc-T6)	NIM OCCIO	ADNTI 2	97
galactosamine:polypeptide N- acetylgalactosaminyltransferase 6 (Gal-	NW 001210	GALIVIO	
UDP-N-acetyl-alpha-D-	NM 007210	GALNT6	96
keratin 23 (histone deacetylase inducible)	MW_0 135 15,	IXI 23	
peptide	NM 014314 NM 015515.	RIG-I KRT23	95
DEAD/H (Asp-Glu-Ala-Asp/His) box poly-		DIO 1	94
like lung adenocarcinoma			
hypothetical protein FLJ20647 membrane protein expressed in epithelial-	NM 024792	CT120	93
hypothetical protein FLJ20315	NM 017763 NM 017918	FLJ20315 FLJ20647	92
sestrin 1	NM_014454	SESN1 FLJ20315	90 91
gen 112	NINA 044454	CECNIA	90
hepatocellular carcinoma-associated anti-	NM_018487	HCA112	89
	NM_080796		121
death associated transcription factor 1	NM_022105,	DATF1	00
CGI-85 protein	NM_016028	CGI-85	87 88
	AUL 4 04 0000	001.05	87

10. The method of claims 1, 2 or 3, wherein at least two of said plurality of gene expression products forming a pattern used to determine said microsatellite status are selected individually from the group consisting of the genes listed below

Gene name	Ref seq	Gene symbol	SEQ NO.:	ID
heterogeneous nuclear ribonucle oprotein L metastais-associated gene family, member 2 chemokine (C-X-C motif) ligand 10 splicing factor, arginine/serine-rich 6	NM 001533 NM 004739 NM 001565 NM 006275	HNRPL MTA2 CXCL10 SFRS6	11 23 35 43	

protein kinase C binding protein 1	NM_012408 NM_183047	PRKCBP1	57 124
hepatocellular carcinoma-associated antigen	NM 018487	HCA112	89
hypothetical protein FLJ20618 SET translocation (myeloid leukaemia- associated)	NM 017903 NM 003011.1	FLJ20618 SET	102 103
ATPase, class II, type 9a	Xm 030577.9	ATP9a	104

11. The method of claims 1, 2 or 3, wherein at least two of said plurality of gene expression products forming a pattern used to determine said microsatellite status are selected individually from the group consisting of the genes listed below

Gene name	Ref seq	Gene symbol	SEQ NO.:	ID
heterogeneous nuclear ribonucleoprotein L	NM 001533	HNRPL	11	
metastais-associated gene family, member 2	NM 004739	MTA2	23	
chemokine (C-X-C motif) ligand 10	NM 001565	CXCL10	35	
splicing factor, arginine/serine-rich 6	NM 006275	SFRS6	43	
protein kinase C binding protein 1	NM_012408	PRKCBP1	57	
hepatocellular carcinoma-associated antigen 112	NM_183047 NM_018487	HCA112	124 89	
hypothetical protein FLJ20618	NM 017903	FLJ20618	102	
SET translocation (myeloid leukaemia- associated)	NM_003011. 1	SET	103	
ATPase, class II, type 9a	Xm_030577. 9	ATP9a	104	

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12. The method of claims 1, 2 or 3, wherein

 i) at least one of said plurality of gene expression products forming a pattern used to determine said microsatellite status is selected from the group of genes consisting of

Gene name	Ref seq	Gene symbol	SEQ NO.:	ID
heterogeneous nuclear ribonucleoprotein L metastais-associated gene family, member		HNRPL	11 23	
2	NM_004739	MTA2	23	

 Chemokine (C-X-C motif) ligand 10
 NM 001565
 CXCL10
 35

 splicing factor, arginine/serine-rich 6
 NM 006275
 SFRS6
 43

and

ii) at least one of said plurality of gene expression products forming a pattern used to determine said microsatellite status is selected from the group of genes consisting of

Gene name	Ref seq	Gene symbol	SEQ ID NO.:
protein kinase C binding protein 1	NM_012408 NM_183047	PRKCBP1	57 124
hepatocellular carcinoma-associated anti- gen 112	NM_018487	HCA112	89
hypothetical protein FLJ20618 SET translocation (myeloid leukaemia- associated)	NM 017903 NM 003011.1	FLJ20618 SET	102 103
ATPase, class II, type 9a	Xm 030577.9	ATP9a	104

13. The method of claims 1, 2 or 3, wherein

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i) at least one of said plurality of gene expression products forming a pattern used to determine said microsatellite status is selected from the group of genes that are down regulated in MSS colon cancers compared to MSI colon cancers consisting of

	•		
Gene name	Ref seq	Gene symbol	SEQ ID NO.:
heterogeneous nuclear ribonucleoprotein L	NM 001533	HNRPL	11
metastais-associated gene family, member 2	NM 004739	MTA2	23
chemokine (C-X-C motif) ligand 10 Splicing factor, arginine/serine-rich 6	NM 001565 NM 006275	CXCL10 SFRS6	35 43

and

ii) at least one of said plurality of gene expression products forming a pattern used to determine said microsatellite status is selected from

the group of genes that are up regulated in MSS colon cancers compared to MSI colon cancers consisting of

Gene name	Ref seq	Gene symbol	SEQ ID NO.:
protein kinase C binding protein 1	NM_012408 NM_183047	PRKCBP1	57 124
hepatocellular carcinoma-associated	NM 018487	HCA112	89
antigen 112 hypothetical protein FLJ20618 SET translocation (myeloid leukaemia- associated)	NM 017903 NM 003011.1	FLJ20618 SET	102 103
ATPase, class II, type 9a	Xm 030577.9	ATP9a	104

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- 14. The method of claim 13, wherein the difference in the level of the gene expression products forming a pattern is at least one-fold.
- 15. The method of claim 13, wherein the difference of the level of the gene expression products forming a pattern is at least 1.5 fold.

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16. The method of claim 1, 2 or 3, wherein at least one of said plurality of gene expression products used to determine the hereditary or sporadic nature of said colon cancer are selected individually from the group consisting of the genes as listed below

Gene name	Ref seq	Gene symbol	SEQ ID NO.:
Homeo box C6	NM_004503	HOXC6	105
Piwi – like 1	NM_004764.2	2PIWIL1	106 .
Mut L homolog 1	NM 00249.2	MLH1	107
Collapsin response mediator protein 1	NM 001313.2	CRMP1	108
Homeo box B2	NM 002145.2	2HOXB2	109
TIOTHEO BOX B2	NM 002860.2	PYCS/ADH18	3 1 1 0
Pyrroline-5-carboxylate synthetase (glutarnate gamma-semialdehyd synthetase		A1	
TGFB inducible early growth response	NM 005655.1	TIEG	111
Checkpoint with forkhead and ring fingedomains??	erNM_018223.	CHFR	112
Hypothetical protein FLJ13842	NM 024645.	1 FLJ13842	113
Phosphoprotein regulated by mitogen pathways			114

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17. The method of claim 1, 2 or 3, wherein at least two of said plurality of gene expression products forming a pattern used to determine said hereditary or sporadic nature of colon cancer are the two genes as listed below

Gene name	Ref seq Gene symbol SEQ ID I NO.:	
Piwi – like 1	NM_004764.2PIWIL1 106	
Mut L homolog 1	NM_00249.2 MLH1 107	

18. The method according to claims 1, 2 or 3, wherein the microsatellite status in an individual having contracted colon cancer is microsatellite instable.

19. The method according to any of the preceding claims, wherein said colon cancer is of Duke's B or Duke's C stage.

- 20. The method according to any of the preceding claims, wherein said colon cancer is an adenocarcinoma, a carcinoma, a teratoma, a sarcoma, and/or a lymphoma.
- 21. The method according to any of the preceding claims, wherein the sample is a biopsy of the tissue.
- 22. The method according to any of the preceding claims, wherein the sample is a cell suspension made from the tissue.
- 23. The method according to any of the preceding claims, wherein the expression level is determined by determining mRNA of the sample.
- 24. The method according to any of the preceding claims, wherein the expression level is determined by determining expression products, such as peptides and/or protein in the sample.

25. The method according to any of the preceding claims, wherein the microsatellite status of the colon cancer in an individual has been determined prior to the determination of the presence and/or amount of gene expression products

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26. The method according to any of the preceding claims, wherein the sporadic or hereditary nature of a colon cancer has been determined prior to the determination of the presence and/or amount of gene expression

products.

- 27. A method for classification of cancer in an individual having contracted cancer, wherein the microsatellite status is determined by a method comprising the steps of
 - i) in a sample from the individual having contracted cancer, said sample comprising a plurality of gene expression products the presence and/or amount of which forms a pattern that is indicative of the microsatellite status of said cancer,
 - ii) determining the presence and/or amount of said gene expression products forming said pattern,
 - iii) obtaining an indication of the microsatellite status of said cancer in the individual based on step ii).
- 28. A method for classification of cancer in an individual having contracted cancer, wherein the hereditary or sporadic nature of the cancer is determined by a method comprising the steps of
 - i) in a sample in a sample from the individual having contracted cancer, said sample comprising a plurality of gene expression products the presence and/or amount of which forms a pattern that is indicative of the hereditary or sporadic nature of said cancer,
 - ii) determining the presence and/or amount of said gene expression products forming said pattern,
 - iii) obtaining an indication of the hereditary or sporadic nature of said cancer in the individual based on step ii).

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29. The method according to claim 28, wherein the microsatellite status of said cancer is determined simultaneously or sequentially therewith. 30. A method for treatment of an individual comprising the steps of 5 i) selecting an individual having contracted a colon cancer, wherein the microsatellite status is stable, determined according to the method of claims 1, 2, 3, 27 or 28 ii) treating the individual with anti cancer drugs 10 31. The method of treatment according to claim 30, wherein the anti cancer drug is selected from the group of fluorouracil-based drugs. 32. The method of treatment according to claim 31, wherein the anti cancer drug is selected from 5-fluorouracil, N-methy-N'-nitro-N-nitro soguanidine 15 and/or 6-thioguanine. 33. The method of treatment according to claim 30, wherein the anti cancer drug is selected from the group of non-fluorouracil based drugs. 20 34. The method according to claim 33, wherein the anti cancer drug is selected from leucovorin, irrinotecan, oxaliplatin, cetuximab. 35. A method for treatment of an individual comprising the steps of i) selecting an individual having contracted a colon can cer, wherein 25 the microsatellite status is instable, determined according to the method of claims 1, 2, 3, 27 or 34 ii) treating the individual with anti cancer drugs. 36. The method according to claim 35, wherein the anti cancer drug is se-30 lected from campothecin or irinotecan.

37. The method according to claim 30 or 35, wherein the microsatellite status has been determined by microsatellite analysis, ELISA, antibody-based

histochemical staining, immuno histo chemistry.

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- 38. The method according to claim 30 or 35 wherein the sporadic or hereditary nature of colon cancer has been examined prior to determining the sporadic or hereditary nature of colon cancer by gene expression products forming a pattern.
- 39. The method according to claim 30 or 35 wherein the sporactic or hereditary nature of colon cancer has been examined by histological examination of the sample.
- 40. The method according to claim 30 or 35 wherein the sporadic or hereditary nature of colon cancer has been examined by genotyping the sample.
- 41. A method for reducing malignancy of a cell, said method comprising contacting a tumor cell in question with at least one pepticle expressed by at least one gene selected from genes being expressed in an at least two-fold higher in tumor cells than the amount expressed in said tumor cell in question.

42. The method according to claim 41, wherein the at least one peptide is selected individually from genes comprising a sequence as identified below

Gene name	Ref seq	Gene symbol	SEQ NO.:	ID
heterogeneous nuclear ribonucleopro- tein L metastais-associated gene family, member 2 chemokine (C-X-C motif) ligand 10	NM 001533 NM 004739 NM 001565 NM 006275	HNRPL MTA2 CXCL10 SFRS6	11 23 35 43	

43. The method according to claim 41, wherein the at least one peptide is selected individually from genes comprising a sequence as identified below

Gene name	Ref seq	Gene	SEQ ID

		symbol	NO.:
protein kinase C binding protein 1	NM_012408 NM_183047	PRKCBP1	57 124
hepatocellular carcinoma-associated	NM_018487	HCA112	89
antigen 112 hypothetical protein FLJ20618 SET translocation (myeloid leukaemia- associated)	NM_017903 NM_003011.1	FLJ20618 SET	102 103
ATPase, class II, type 9a	Xm_030577.9	ATP9a	104

- 44. The method according to claim 41 or 42, wherein the tumor cell is contacted with at least two different peptides.
- 45. A method for reducing malignancy of a tumor cell in question comprising,
 - i) obtaining at least one gene selected from genes being expressed in at least one fold higher in tumor cells than the amount expressed in the tumor cell in question,
 - ii) introducing said at least one gene into the turnor cell in question in a manner allowing expression of said gene(s).

46. The method according to claim 45, wherein the at least one gene is selected from genes comprising a sequence as identified below

Gene name	Ref seq	Gene symbol	SEQ NO.:	ID
Heterogeneous nuclear ribonucleoprotein L metastais-associated gene family, member	NM 001533	HŅRPL	11 23	
2 Chemokine (C-X-C motif) ligand 10 splicing factor, arginine/serine-rich 6	NM 004739 NM 001565 NM 006275	MTA2 CXCL10 SFRS6	35 43	

47. The method according to claim 45, wherein the at least one gene is selected from genes comprising a sequence as identified below

Gene name	Ref seq	Gene symbol	SEQ NO.:	ID
Protein kinase C binding protein 1	NM_012408 NM_183047	PRKCBP1	57 129	
hepatocellular carcinoma-associated anti-	NM_018487	HCA112	89	
gen 112 hypothetical protein FLJ20618	NM_017903	FLJ20618	102	

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SET translocation (myeloid leukaemia- associated)	NM_003011.1	SET	103	
ATPase, class II, type 9a	Xm 030577.9	ATP9a	104	

- 48. The method according to claim 45, 46 or 47, wherein at least two different genes are introduced into the tumor cell.
- 49. A method for reducing malignancy of a cell in question, said method comprising
- obtaining at least one nucleotide probe capable of hybridising with at
 least one gene of a tumor cell in question, said at least one gene being
 selected from genes being expressed in an amount at least one-fold
 lower in tumor cells than the amount expressed in said tumor cell in
 question, and
 - introducing said at least one nucleotide probe into the tumor cell in question in a manner allowing the probe to hybridise to the at least one gene, thereby inhibiting expression of said at least one gene.
 - 50. The method according to claim 49, wherein the nucleotide probe is selected from probes capable of hybridising to a nucleotide sequence comprising a sequence as identified below

	Ref sea	Gene	SEQ	ID
Gene name	Nei seq	symbol	NO.:	
protein kinase C binding protein 1	NM_012408 NM_183047	PRKCBP1	57 124	
hepatocellular carcinoma-associated antigen	NM_018487	HCA112	89	
hypothetical protein FLJ20618 SET translocation (myeloid leukaemia- associated)	NM 017903 NM 003011.1	FLJ20618 SET	102 103	
ATPase, class II, type 9a	Xm 030577.9	ATP9a	104	

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51. The method according to claim 46, wherein the nucleotide probe is selected from probes capable of hybridising to a nucleotide sequence comprising a sequence as identified below

Gene name	Ref seq	Gene symbol	SEQ NO.:	ΙĎ
heterogeneous nuclear ribonucleoprotein L metastais-associated gene family, member 2 chemokine (C-X-C motif) ligand 10 splicing factor, arginine/serine-rich 6	NM 001533 NM 004739 NM 001565 NM 006275	HNRPL MTA2 CXCL10 SFRS6	11 23 35 43	

52. The method according to claim 49, 50 or 51, wherein at least two different probes are introduced into the tumor cell.

53. A method for producing antibodies against an expression product of a cell from a biological tissue, said method comprising the steps of

obtaining expression product(s) from at least one gene said gene being expressed as defined in any of claims 1-29,

immunising a mammal with said expression product(s) obtaining antibodies against the expression product.

- 54. A method for treatment of an individual comprising the steps of
 - i) selecting an individual having contracted a colon cancer, wherein the microsatellite status is stable, determined according to the method of claims 1, 2, 3, 27 or 28 and wherein the hereditary nature of said cancer has been determined according to the method of claims 1, 2 or 3
 - ii) introducing at least one gene into the tumor cell in a manner allowing expression of said gene(s).
- 55. The method according to claim 54, wherein the at least one gene is selected from MSH2, MLH1, PMS1, PMS2 or MSH6.

Gene name	Ref seq	Gene symbol	SEQ NO.:	ID
Homo sapiens mutS homolog 2, colon cancer, nonpolyposis type 1 (E. coli)	NM_000251	MSH2	136	
Mut L homolog 1 Homo sapiens PMS1 postmeiotic segregation increased 1 (S. cerevisiae)	NM_00249.2 NM_000534	MLH1 PMS1	107 137	
Homo sapiens PMS2 postmeiotic segregation	NM_000535	PMS2	138	
increased 2 (S. cerevisiae) (PMS2), mRNA Homo sapiens mutS homolog 6 (E. coli)	NM_000179	MSH6	139	

- 56. The method according to claim 54 or 55, wherein at least two different genes are introduced.
- 57. Pharmaceutical composition for the treatment of a classified cancer comprising at least one antibody as defined in claim 53.
 - 58. Pharmaceutical composition for the treatment of a classified cancer , comprising at least one polypeptide as defined in any of the claims 41-44.
 - 59. Pharmaceutical composition for the treatment of a classified cancer comprising at least one nucleic acid and/or probe as defined in any of the claims 45-52.
 - 60. The use of a method as defined in any of claims 1- 37 for producing an assay for classifying cancer in animal tissue.
 - 61. The use of a peptide as defined in any of claims 41-44 for preparation of a pharmaceutical composition for the treatment of a cancer in animal tissue.
 - 62. The use of a gene as defined in any of claims 45-52 for preparation of a pharmaceutical composition for the treatment of cancer in animal tissue.

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63. The use of a probe as defined in any of claims 49-52 for preparation of a pharmaceutical composition for the treatment of cancer in animal tissue. 64. An assay for classification of cancer in an individual having contracted 5 cancer, comprising at least one marker capable of determining the microsatellite status in a sample and at least one marker in a sample determining the prognostic marker, 10 wherein the microsatellite status and the prognostic marker is determined simultaneously or sequentially. 65. The assay according to claim 64, wherein the marker is a nucleotide probe. 15 66. The assay according to claim 64, wherein the marker is an antibody. 67. The assay according to claim 64, wherein the genes are as defined in 20 any of claims 9-13 or 16-17.